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(54) Title: AMYLIN AND POSSIBLY INSULIN CONTAINING COMPOSITION FOR THE TREATMENT OF ANOR-EXIA AND RELATED STATES

(57) Abstract

Method for treatment of a patient suffering from anorexia or a related condition by administering amylin or an analogue thereof to the patient in an amount sufficient to increase amylin and/or insulin levels in the plasma of the patient.

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AMYLIN AND POSSIBLY INSULIN CONTAINING COMPOSITION FOR THE TREATMENT OF ANOREXIA AND RELATED STATES

DESCRIPTION

Treatment of Anorexia and Related States

Related Applications

This application is continuation-in-part of U.S. Serial No. 07/862,500 filed April 3, 1992, and a continuation-in-part of U.S. Serial No. 07/704,995, filed May 24, 1991, the contents of which are both hereby incorporated by reference.

Background of the Invention

This invention relates to treatment of anorexia and related states.

Anorexia, defined as the lack or the loss of appetite for food (Dorland's Illustrated Medical Dictionary, 24 edition, W.B. Saunders Company, Philadelphia, 1965) has multiple etiologies. It is commonly associated with cachexia, "a profound and marked state of constitutional disorder, general ill health and malnutrition" [idem]. Common examples of conditions associated with anorexia and cachexia are anorexia nervosa, certain infectious diseases, and malignancy.

Anorexia nervosa is a serious psychiatric disorder 20 affecting predominantly women (94-96%) in the 13-30 age range. Between 1% (Crisp et al., 128 Br. J. Psychiatry 549, 1976) and 3% (Ballot et al., 59 S.Afr. Med. J. 992, 1981) of young women may be affected. The morbidity and mortality from this condition are considerable. Two years 25 from diagnosis, 4-6% have died and only 50% have achieved a normal weight. There are multiple endocrine and metabolic abnormalities present, most of which are believed to be secondary to the malnutrition. A serious complication of the condition is osteoporosis, which can involve both 30 the spine and peripheral bones. At present there is no specific treatment for anorexia nervosa, although multiple approaches have been tried (Piazza, Piazza & Rollins Compr. Psychiatry 21:177-189 1980).

In experimental animals, infection with various agents such as Mycobacterium bovis (Carswell et al., Proc. Natl. Acad. Sci. U.S.A. 73:3666-3670 1975) caused the appearance of a blood factor that caused necrosis of 5 tumors in mice (tumor necrosis factor, TNF). In a different line of investigation, an agent produced in response to infection with Trypanosoma brucei produced unexpected weight loss an wasting (Rouzer & Cerami Mol. Biochem. Parasitol 2:31-38 1980), and was termed cachexin. TNF and 10 cachexin have since been shown to be the same 17kD protein, produced by activated macrophages. It stimulates several aspects of the immune response. Injected into animals, it produces many of the features of cachexia, including anorexia, bone resorption, and the inhibition of 15 fat uptake into adipocytes. It has been proposed that this agent, which might be produced by tumor cells, or as a host response, could account for some of the cachexia of cancer (Cerami et al., Recent Prog. Horm. Res. 43:99-112 However, the association of cachexin/TNF with 20 malignancy is the subject of conflicting reports.

Nutritional support via either enteral (via the gut) or parenteral (e.g., intravenous) therapy is indicated in patients unable to take sufficient nutrition by mouth, and who are therefore at risk for the complication of malnutrition. Therapy attempts to maintain anabolism (buildup of body substance stores) and avert catabolism (breakdown). To this end, insulin is commonly added to intravenously infused nutrients. Examples of patients requiring parenteral nutrition or other nutritional support include those with inflammatory bowel diseases, patients with resected bowel, severe preoperative malnutrition and acute pancreatitis (Howard, In: Harrison's Principals of Internal Medicine 12th Edition, Wilson et al. (eds), McGraw-Hill, New York 1991, p. 429).

35 Human diabetics are deficient in insulin secretion, and in some cases lack insulin. Insulin is one of several hormones which play a role in regulation of blood glucose

levels. Simplistically, there are two main stores of glucose in a mammal—the liver and skeletal muscle, where glucose is stored in the form of glycogen. Muscle glycogen is used as a glucose source for the muscle, whereas liver glycogen is used as a glucose source for all tissues, including blood. It is the interplay of certain hormones in regulation of glycogen accumulation and breakdown that is critical in the invention described below.

Insulin regulates glucose uptake by muscle tissue for storage of the glucose as muscle glycogen. Insulin also prevents hyperglycemia, that is, the unacceptable accumulation of high levels of glucose in the blood, and suppresses conversion of liver glycogen to glucose, and subsequent secretion of that glucose into the blood. In the presence of excess insulin, blood glucose accumulates in muscle tissue as glycogen, liver glucose output is suppressed, and the level of blood glucose falls, to create a condition termed hypoglycemia.

Another hormone, glucagon, increases blood glucose levels by stimulating liver glycogen breakdown to glucose, and subsequent secretion of that glucose. This liver glycogen is used to maintain blood glucose levels, and glucagon may be considered an insulin counterregulatory hormone.

Amylin is another hormone which has been discovered to be concerned in regulation of blood glucose levels. It reverses insulin-mediated suppression of liver glucose output in rats. Molina et al., 39 Diabetes 260, 1990, and Koopmans et al., 39 Diabetes 101A, 1990.

European Patent Application No. 88307927.9 describes the treatment of diabetes mellitus or hypoglycemia with amylin, or with a combination of amylin and insulin, preferably at a ratio of between 100:1 to 0.1:1 insulin to amylin.

Summary of the Invention

Applicant have discovered that a patient suffering from anorexia may have fasting plasma amylin and insulin concentrations below the normal range, and in fact near 5 the range measured by Type 1 diabetics. Applicant believe that patients suffering from cachexia or receiving parenteral nutrition (i.e., nutrition except oral nutrition, e.g., intravenous) have reduced amylin and/or insulin levels. Thus, applicants propose that patients suf-10 fering from anorexia and cachectic states, as well as patients undergoing parenteral nutrition, be administered amylin with or without insulin. Such administration will preferably increase adipose tissue in such patients and thus be of significant benefit. The amount of the hor-15 mones (amylin or amylin and insulin) that are administered should preferably be sufficient to increase the hormone plasma levels of the patient to normal levels observed in the general population. For example, in a patient having a level of insulin similar to that in a Type 1 diabetic, it is necessary to administer about 1 mg per day of amylin, alone or with insulin together in a ratio of amylin to insulin between 1:100 and 10:1.

Thus, in a first aspect the invention features a method for treatment of a patient suffering from anorexia by administering amylin or an agonist analogue thereof to the patient in an amount sufficient to increase the amylin level in the plasma of the patient.

In related aspects, the invention features methods for treatment of cachectic patients and those undergoing parenteral nutrition by similarly administering amylin or an agonist analogue thereof in an amount sufficient to increase the amylin plasma level.

In preferred embodiments of the above aspects, the invention features co-administering insulin to the patient in an amount sufficient to increase insulin plasma level; and the amylin (or agonist analogue) and insulin are provided in an amount sufficient to increase the level of

adipose tissue in the patient. For example, the amount of amylin or agonist analogue provided is sufficient to increase liver glycogen stores. Insulin and amylin in the patient will act together to enhance the deposition of body fat. Insulin will enhance the uptake of glucose into fat cells and will enhance the transfer of lipid into fats cells via activation of lipoprotein lipase at adipose tissue capillaries. Amylin will enhance the hepatic supply of lactate, a favored lipogenic substrate (Carmona & Freedland 119 J. Nutr. 1304, 1989).

In another related aspect, the invention features a method for treating a patient that is deficient in adipose tissue by administering amylin (or an agonist analogue thereof) and/or insulin as described above in an amount sufficient to increase the amount of adipose tissue in that patient. Those in the art will recognize that standard procedures can be used to measure the increase in such adipose tissues, and to identify those patients which are deficient in adipose tissue levels.

In preferred embodiments, the method includes the step of identifying a mammal having the above-noted conditions, prior to the administering step.

In other preferred embodiments, combinations of amylin and insulin are provided in a molar ratio between 25 about 1:2.5 and 1:35 or about 1:5 and 1:25, and at least 0.5 micrograms of amylin per kilogram of the patient per day are provided.

The level of insulin and amylin in the patient may be determined by any desired means, many examples of which 30 exist in the published literature. In addition, amylin activity can be assayed as described by Cooper and Young, U.S. Serial No. 07/666,512, entitled "Amylin Activity Assays", filed March 8, 1991, assigned to the same assignee as the present application, and hereby incorporated by reference herein.

Preferably, where amylin and insulin are administered rather than amylin alone, the composition includes an

amylin and an insulin in a molar ratio of between about 1:2.5 and 1:35, preferably in a form which allows delayed release of both the insulin and amylin in a constant molar ratio, or in a form suitable for parenteral administration.

In a related aspect, the treatment may include administering a composition containing an insulin and an amylin at a suitable molar ratio, such that the amount of amylin in the composition will result in circulating plasma levels of amylin that are about 3 to about 6% that of insulin upon administration of the composition to the patient.

The term "amylin" is used in this application to include compounds defined by Young et al., U.S. applica-15 tion Serial No. 07/640,478, filed January 10, entitled "Hyperglycemic Compositions", which (including drawings) is hereby incorporated by reference. For example, it includes the peptide hormone, and species variations of it, referred to as amylin which is synthesized 20 and secreted from the beta cells of the pancreas. it includes human amylin, cat amylin, dog amylin, rat amylin, mouse amylin, hamster amylin, and guinea pig Preferably the amylin has an EC50 of less than 1 nM in the rat soleus muscle assay as described in, for 25 example, European Patent Application 88307927.9. functions along with insulin, which is stored and released from the same pancreatic beta cells, to regulate fuel Amylin acts through receptors located in metabolism. skeletal muscle to increase glycogen turnover in this 30 tissue, believed to result in an increased return to the bloodstream of lactate, which is a major precursor of hepatic gluconeogenesis. Amylin cosecretion with insulin after meals therefore results in restoration of hepatic glycogen content and limits the potential which would 35 otherwise exist for insulin to induce hypoglycemia. Administration of amylin to anesthetized rats produces large increases in blood lactate levels, presumably

through a direct effect upon skeletal muscle glycogen breakdown and glycolysis. Increased blood lactate content is followed rapidly by increased blood glucose levels, believed to result from provision of gluconeogenic precursors in the form of lactate to the liver. These physiological and pharmacological effects of amylin form the basis for its therapeutic indications in treatment of Type 1 diabetes and hypoglycemia.

The term "amylin analogue" or "agonist analogue"

includes derivatives of amylin or other reagents aching as
agonists at an amylin receptor, or having those biological
properties described above.

The term "analogue" is meant to include human amylin equivalents known to those of ordinary skill in the art.

15 For example, various amino acids in the amylin sequence can be substituted with equivalent amino acids in a manner which has little (i.e., reduces activity less than 20%) or no effect on the biological activity of the amylin, as measured in the assay described above. For example, neutral amino acids, and charged amino acids replaced with equivalently charged amino acids. In addition, one or more amino acids may be deleted from the polypeptide if such deletion has little or no affect on the biological activity of the human amylin.

Useful analogues include agonist analogues identified in copending and commonly assigned U.S. Serial No. 07/794,266 filed November 19, 1991, the contents of which are hereby incorporated by reference. In particular, useful analogues include agonist analogues having the following sequence:

 $^{1}A_{i}-X-Asn-Thr-^{5}Ala-Thr-Y-Ala-Thr-^{10}Gln-Arg-Leu-B_{i}-Asn-^{15}Phe-Leu-C_{i}-D_{i}-E_{i}-^{20}F_{i}-G_{i}-Asn-H_{i}-Gly-^{25}I_{i}-J_{i}-Leu-K_{i}-I_{i}-^{30}Thr-M_{i}-Val-Gly-Ser-^{35}Asn-Thr-Tyr-Z$

wherein A_1 is hydrogen Lys, Ser, Ala, des- α -amino Lys, or acetylated Lys; B_1 is Ala, Ser or Thr; C_1 is Val, Leu or Ile; D_1 is His or Arg; E_1 is Ser or Thr; F_1 is Ser, Thr, Gln

or Asn; G, is Asn, Gln or His; H, is Phe, Leu or Tyr; I, is Ala or Pro; J1 is Ile, Val, Ala or Leu; K1 is Ser, Pro, Leu, Ile or Thr; Li is Ser, Pro or Thr; Mi is Asn, Asp or Gln; X and Y are independently selected residues having 5 side chains which are chemically bonded to each other to form an intramolecular linkage; and Z is hydroxy, amino, alkylamino, dialkylamino, cycloalkylamino, arylamino, aralkylamino, alkyloxy, aryloxy or aralkyloxy; provided that (a) when A_i is Lys, B_i is Ala, C_i is Val, D_i is His, E_i 10 is Ser, F₁ is Ser, G₁ is Asn, H₁ is Phe, I₁ is Ala, J₁ is Ile, K_i is Ser, L_i is Ser, and M_i is Asn; (b) when A_i is Lys, B_1 is Ala, C_1 is Ile, D_1 is Arg, E_1 is Ser, F_1 is Ser, G, is Asn, H, is Leu, I, is Ala, J, is Ile, K, is Ser, L, is Pro, and M_1 is Asn; (c) when A_1 is Lys, B_1 is Ala, C_1 is 15 Val, D_i is Arg, E_i is Thr, F_i is Ser, G_i is Asn, H_i is Leu, I_i is Ala, J_i is Ile, K_i is Ser, L_i is Pro, and M_i is Asn; (d) when A_1 is Lys, B_1 is Ala, C_1 is Val, D_1 is Arg, E_1 is Ser, F1 is Ser, G1 is Asn, H1 is Leu, I1 is Pro, J1 is Val, K_i is Pro, L_i is Pro, and M_i is Asn; (e) when A_i is Lys, B_i 20 is Ala, C_1 is Val, D_1 is His, E_1 is Ser, F_1 is Asn, G_1 is Asn, H1 is Leu, I1 is Pro, J1 is Val, K1 is Ser, L1 is Pro and M, is Asn; or (f) when A, is Lys, B, is Thr, C, is Val, D_i is Arg, E_i is Ser, F_i is Ser, G_i is His, H_i is Leu, I_i is Ala, J_1 is Ala, K_1 is Leu, L_1 is Pro and M_1 is Asp; then one 25 or more of any of A_1 to M_1 is not an L-amino acid and Z is not amino.

Suitable side chains for X and Y include groups derived from alkyl sulfhydryls which may form disulfide bonds; alkyl acids and alkyl amines which may form cyclic lactams; alkyl aldehydes or alkyl halides and alkylamines which may condense and be reduced to form an alkyl amine bridge; or side chains which may be connected to form an alkyl, alkenyl, alkynyl, ether or thioether bond. Preferred alkyl chains include lower alkyl groups having from about 1 to about 6 carbon atoms.

The term "alkyl" refers to both straight— and branched—chain alkyl groups. The term "lower alkyl" refers to both straight— and branched—chain alkyl groups having a total of from 1 to 6 carbon atoms and includes primary, secondary and tertiary alkyl groups. Typical lower alkyls include, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, isobutyl, t-butyl, n-pentyl, n-hexyl, and the like.

The term "aryl" refers to carbocyclic aromatic groups
of 6 to 14 carbon atoms such as phenyl and naphthyl, as
well as heterocyclic aromatic groups containing 1 to
3 heteroatoms (nitrogen, oxygen, sulfur, etc.) such as
pyridyl, triazolopyrazine, pyrimidine and the like.

The term "aralkyl" refers to an "aryl" group of 6 to 10 carbon atoms directly attached to an "alkyl" group of 1 to 4 carbon atoms and includes for example benzyl, p-chlorobenzyl, p-methylbenzyl, and 2-phenylethyl.

The term "cycloalkyl" refers to cyclic alkyl groups of 5 to 8 carbon atoms.

By "identifying" is meant to include noting the symptoms or characteristics of anorexia or cachectic conditions. Such symptoms are well known in the art. It also includes chemical or biochemical assays which indicate such conditions, or their equivalent.

By "insulin" is meant a polypeptide or its equivalent useful in regulation of blood glucose levels. A general description of such insulins is provided in Goodman and Gilman, "The pharmacological basis of therapeutics", 7th ed., Maxmillan Pub. Co. at e.g., p. 1501, et seq. (1985).

30 Such insulins can be fast acting, intermediate acting, or long acting. <u>Id.</u> at 1502. Various derivatives of insulin exist and are useful in this invention. <u>See</u>, <u>e.g.</u>, U.S. Patents, 5,049,547, 5,028,587, 5,028,586, 5,016,643. Insulin peptides are also useful (<u>see</u>, <u>e.g.</u>, U.S. Patent

5,008,241), as are analogues (see, e.g., U.S. Patent 4,992,417 and 4,992,418). Such insulin can be administered by any standard route, including nasal administra-

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tion, see, e.g., U.S. Patents 4,988,512 and 4,985,242, and 2 Bioworld Today, No. 125, 1, 1991.

Other features and advantages of the invention will be apparent from the following description of the pre-5 ferred embodiments thereof, and from the claims.

Description of the Preferred Embodiments Fat Deficient Conditions

Conditions such as anorexia nervosa, cachectic conditions and the general condition of patients receiving 10 intravenous nutrition or other related conditions may be In each of these conditions treated in this invention. there is either progressive loss of both adipose tissue and lean body mass or failure to increase these from a base of marked established loss.

Applicant have found that amylin is a controller of fuel cycling from muscle to liver and liver to peripheral Without being bound to any particular theory, Applicant believes that this is due, at least in some part, to promotion of Cori cycling of carbohydrate from 20 skeletal muscle glycogen to liver glycogen and by the provision of hepatic substrate for triglyceride synthesis. Applicant believes that activation of these pathways improves the efficient storage of food-derived substrates in liver and in adipose tissue in the above-exemplified 25 conditions. Thus, the combined use of amylin and insulin in which amylin serves to provide hepatic substrate and insulin promotes hepatic production of triglyceride and lipogenesis in adipose tissue is beneficial.

Promotion of formation of adipose tissue is critical 30 to normal health not only as a concentrated store of energy for use in fasting or exercise, but subcutaneous fat especially is important in establishing the body contours and cushioning of the underlying tissues. sores, for example, may be caused or exacerbated by loss 35 of adipose tissue.

Amylin deficiency in Type 1 diabetics has been proposed as a pathologic basis for difficulties in achieving good glycemic control with insulin therapy. Applicant discovered that an anorexic patient has fasting plasma amylin and insulin concentrations below the normal range, and in fact near the range measured in Type 1 diabetics. When such a patient was administered a standard 75 g glucose oral load there was a very small transient deviation of amylin and insulin levels, markedly lower than that seen in normal subjects. Thus, it appears that anorexia nervosa is an amylin, and possibly insulin, deficient state which can be treated by administration of amylin and/or insulin.

While this proposal is counter to reports that amylin can suppress appetite (which is clearly an undesirable feature for treatment of anorexia or cachectic states), applicant believes that the appetite suppressant effects of amylin is seen only at very high doses and may be short lived. Indeed, applicant has discovered that in toxicological studies with amylin in both rats and dogs, where two weeks of amylin administration were used, there was no reduction in food intake or weight in the animal.

As with anorexia, applicant believes that both cachectic states and patients receiving total parenteral nutrition are amylin and/or insulin deficient states and thus, appropriate for amylin replacement or augmentation therapy.

Compositions

Compositions or products according to the invention may conveniently be provided in the form of solutions suitable for parenteral or nasal or oral administration. In many cases, it will be convenient to provide an amylin or insulin in a single solution for administration together. In other cases, it may be more advantageous to administer amylin and insulin separately. A suitable administration regime may best be determined by a doctor

for each patient individually. It will generally be preferable to formulate such that the molar ratio of amylin and insulin for the treatment is between 1:100 and 10:1, or between 1:2.5 and 1:35. Most preferably between 5 1:25 or 1:20 and 1:5.

Since the products of the invention are amphoteric they may be utilized as free bases, as acid addition salts or as metal salts. The salts must, of course, be pharmaceutically acceptable, and these will include metal salts, particularly alkali and alkaline earth metal salts, e.g., potassium or sodium salts. A wide variety of pharmaceutically acceptable acid addition salts are available. These include those prepared from both organic and inorganic acids, preferably mineral acids. Typical acids which may be mentioned by way of example include citric, succinic, lactic, hydrochloric and hydrobromic acids. Such products are readily prepared by procedures well known to those skilled in the art.

The products of the invention will normally be provided as parenteral compositions for injection or infusion. They can, for example, be suspended in an inert
oil, suitably a vegetable oil such as sesame, peanut, or
olive oil. Alternatively, they can be suspended in an
aqueous isotonic buffer solution at a pH of about 5.6 to
7.4. Useful buffers include sodium citrate-citric acid
and sodium phosphate-phosphoric acid.

The desired isotonicity may be accomplished using sodium chloride or other pharmaceutically acceptable agents such as dextrose, boric acid, sodium tartrate, propylene glycol or other inorganic or organic solutes. Sodium chloride is preferred particularly for buffers containing sodium ions.

If desired, solutions of the above compositions may be thickened with a thickening agent such as methyl cellulose. They may be prepared in emulsified form, either water in oil or oil in water. Any of a wide variety of pharmaceutically acceptable emulsifying agents may be

employed including, for example acacia powder, or an alkali polyether alcohol sulfate or sulfonate such as a Triton.

The therapeutically useful compositions of the invention are prepared by mixing the ingredients following generally accepted procedures. For example, the selected
components may be simply mixed in a blender or other standard device to produce a concentrated mixture which may
then be adjusted to the final concentration and viscosity
by the addition of water or thickening agent and possibly
a buffer to control pH or an additional solute to control
tonicity.

For use by the physician, the compositions will be provided in dosage unit form containing an amount of amylin and/or insulin which will be effective in one or multiple doses to control adipose tissue formation at the selected level. As will be recognized by those in the field, an effective amount of therapeutic agent will vary with many factors including the age and weight of the patient, the patient's physical condition, the blood sugar level to be obtained, and other factors. Typical dosage units for treatment of anorexia and related conditions will contain, for example, from about 0.1 to 10 mg of an amylin and about 0.1 to about 1.0 mg of an insulin.

25 Methods

As defined above, compositions useful in the invention are formulated by standard procedure. These compositions are also administered by standard procedure. Suitable doses are readily determined by those in the art, examples of which are provided above.

Amylin analogues may be prepared by using certain conventional coupling reactions known in the peptide art. The analogues are prepared by successively adding the desired amino acid to a growing peptide chain. Typically, an α-N-carbamoyl protected amino acid and an amino acid attached to the growing peptide chain on a resin support

are reacted at room temperature in an inert solvent such as N-methylpyrrolidone, dimethylformamide or methylene chloride in the presence of coupling agents such as dicyclohexylcarbodiimide 1-hydroxybenzotriazole in the 5 presence of a base such as diisopropylethylamine. The α -N-carbamoyl protecting group is removed from the resultant peptide with a reagent such as trifluoroacetic acid or piperidine, and the coupling reaction repeated with the next desired N-protected amino acid. Suitable N-protect-10 ing groups are known in the art, with t-butyloxycarbonyl herein preferred.

Certain preferred methods for synthesis are described in the commonly-assigned copending and commonly assigned patent application Serial No. 667,040 ("Synthetic Prepara-15 tion of Amylin and Amylin Analogs", filed March 8, 1991). These methods provide for solid phase synthesis of a peptide which comprises amylin or an amylin analogue which has enhanced biological activity and is substantially free of deletion and other contaminating peptides wherein said 20 peptide is synthesized using successive synthesis cycles, whereby in each such synthesis cycle, a designated amino acid is added to a growing peptide chain attached to an insoluble resin support by formation of a peptide linkage between an α -amino group of the growing peptide chain and 25 on α -carboxyl of the designated amino acid; and wherein each synthesis cycle comprises: (a) treating the growing peptide chain under α -amino deprotecting conditions to remove an α -amino group; (b) activating the α -carboxyl group of the α -amino protected designated amino acid; (c) contacting the growing peptide chain and the designated amino acid under coupling conditions to form a peptide linkage between the free α -amino for the peptide chain and the activated α -carboxyl of the designated amino acid; and (d) repeating steps (b) and (c) if the coupling 35 efficiency of step (c) is less than about 97%. It is preferred to repeat steps (b) and (c) if the coupling efficiency is less than about 99%. In another preferred

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aspect, steps (b) and (c) are repeated in each synthesis cycle. Optionally, the coupling efficiency is measured after each coupling step.

Suitable coupling conditions include use of a solvent 5 system which maximizes swelling of the solid support, minimizes secondary structure elements of the peptide chain during synthesis cycles, and minimizes intrapeptide and interpeptide hydrogen bonding. Preferably the synthesis cycle includes a capping step after the coupling 10 step(s) wherein unreacted μ -amino groups of the peptide chain are rendered unreactive. The synthesis cycle is successively repeated using appropriate protected α-amino acids to give amylin or an amylin analogue of specified sequence. After completions of the successive synthesis 15 cycles, said amylin or amylin analogue is cleaved from the solid support. It is preferred that the cysteine residues of the peptide chain are selectively deprotected and an intramolecular disulfide bond is formed before cleaving the peptide bond from the solid support.

Suitable α-amino protective groups include t-butoxy-carbonyl and 9-fluorenylmethoxycarbonyl. In one preferred aspect, when t-butoxycarbonyl is used as the α-amino protecting group, the α-carboxyl groups are activated using dicyclohexylcarbodiimide and 1-hydroxybenzotriazole to form 1-hydroxybenzotriazole esters. A particularly preferred solvent system comprise N-methylpyrrolidone.

Amylins and amylin analogues may also be prepared using recombinant DNA techniques, using methods now known in the art. See, e.g., Sambrook et al., Molecular Clon
ing: A Laboratory Manual, 2d Ed., Cold Spring Harbor (1989).

Analogues

Analogues of amylin can be assayed for activity in the soleus muscle assay described above. Amylin agonist activity of compounds may also be assessed by the ability to induce hyperlactemia and/or hyperglycemia in mammals.

The preferred analogues des-Lys-h-amylin, 28Pro-h-amylin, ^{25,28,29}Pro-h-amylin, ¹⁸Arg^{25,28}Pro-h-amylin, and Lys 18 Arg 25,28 Pro-h-amylin, all show amylin activity in vivo in treated test animals, provoking marked hyperlactemia 5 followed by hyperglycemia. In addition to having activities characteristic of amylin, certain of the preferred compounds also possess more desirable solubility and stability characteristics when compared to human amylin. These preferred compounds include 25Pro26Val28,29Pro-h-amylin, 25,28,29Pro-h-amylin, and 18Arg25,28Pro-h-amylin.

Compounds described herein which are especially preferred include 18Arg25,28Pro-h-amylin, des-1Lys18Arg25,28Pro-hamylin, 18Arg^{25,28,29}Pro-h-amylin, des-lLys¹⁸Arg^{25,28,29}Pro-h-amylin, ^{25,28,29}Pro-h-amylin, des-¹Lys^{25,28,29}Pro-h-amylin, and ²⁵Pro²⁶Val^{25,28} Still further amylin analogues include: Pro-h-amylin.

²³Leu²⁵Pro²⁶Val^{28,29}Pro-h-amylin; ²³Leu²⁵Pro²⁶Val²⁸Pro-h-amylin; des-1Lys23Leu25Pro26Val28Pro-h-amylin; ¹⁸Arg²³Leu²⁵Pro²⁶Val²⁸Pro-h-amylin; ¹⁸Arg²³Leu^{25,28,29}Pro-h-amylin; 20 ¹⁸Arg²³Leu^{25,28}Pro-h-amylin; 17Ile23Leu25,28,29Pro-h-amylin; 17Ile^{25,28,29}Pro-h-amylin; des-1Lys17Ile23Leu25,28,29Pro-h-amylin; 17Ile18Arg23Leu-h-amylin; 25 17Ile18Arg23Leu26Val29Pro-h-amylin; ¹⁷Ile¹⁸Arg²³Leu²⁵Pro²⁶Val^{25,29}Pro-h-amylin; 13Thr21His23Leu26Ala28Leu29Pro31Asp-h-amylin; ¹³Thr²¹His²³Leu²⁶Ala²⁹Pro³¹Asp-h-amylin; des-1Lys13Thr21His23Leu26Ala28Pro31Asp-h-amylin; 30 13Thr18Arg21His23Leu26Ala29Pro31Asp-h-amylin; 13Thr18Arg21His23Leu28,29Pro31Asp-h-amylin; and 13Thr18Arg21His23Leu25Pro26Ala28,29Pro31Asp-h-amylin. Other embodiments are within the following claims.

Claims

1. A method for treatment of a patient suffering from anorexia comprising the step of:

administering amylin or an analogue thereof to said patient in an amount sufficient to increase the amylin level in the plasma of said patient.

- 2. A method for treatment of a cachectic patient or a patient undergoing parenteral nutrition comprising the step of:
- administering amylin or an analogue thereof to said patient in an amount sufficient to increase the amylin level in the plasma of said patient.
- 3. The method of claim 1 or 2 further comprising administering insulin to said patient in an amount sufficient to increase the insulin level in said plasma.
 - 4. The method of claim 3 wherein said amylin or insulin are provided in an amount sufficient to increase level of adipose tissue in said patient.
- 5. The method of claim 1 or 2 wherein said amylin 20 is provided in an amount sufficient to increase level of adipose tissue in said patient.
- 6. The method of claim 1 wherein said amount of amylin administered is sufficient to increase the anabolic effect of plasma amylin within the liver, and thereby increase glycogen levels by glycogenesis.
 - 7. The method of claim 1 wherein said insulin and amylin are provided in an amount sufficient to enhance mobilization of lactate from muscle to form fat in the liver.

8. A method for treating a patient that is deficient in adipose tissue by administering amylin and/or insulin in an amount and ratio sufficient to increase adipose tissue in said patient.

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 92/04357

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